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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Application of: Yang, Yinong

Serial No.: 10/768,886

Art Unit: 1638

Filed: January 31, 2004

Examiner: Vinod Kumar

For: Mitogen-Activated Protein Kinase
And Methods for Use to Enhance Biotic
And Abiotic Stress Tolerance in Plants

Atty Docket No.: UAF-03-14

**SUPPLEMENTAL DECLARATION OF YINONG YANG, PH.D.
UNDER 37 C.F.R. §1.131**

Mail Stop Amendment
Commissioner for Patents
P.O. Box 1450
Alexandria, Virginia 22313-1450

I, Yinong Yang certify the following:

1. I am the inventor of U.S. Patent Application No. 10/768,886.
2. The data filed with this declaration was generated from work performed in my laboratory by me or under my direct supervision at the University of Arkansas located in Fayetteville, Arkansas.
3. Prior to 2002, I completed my invention as described and claimed in the above referenced application as evidence below.
4. On or about May 2000, my laboratory isolated the gene fragment of OsMAPK5 (plasmid clone #2C12) (see attached Exhibit A, Lab Notebook I at pages 1-2).
5. On or about September 2000, my laboratory isolated the full length gene of OsMAPK5 (plasmid clone #M2) (see attached Exhibit A, Lab Notebook I at pages 3-5).

6. From approximately November 2000 to May 2001, RNA and protein analysis of OsMAPK5a indicating response to biotic and abiotic stresses were performed in my laboratory (see attached Exhibit A, Lab Notebook I at pages 6- 7).
7. On or about November 2000, rice transformation was initiated for over-expression (H series) and suppression (F series) of OsMAPK5.
8. On or about May 2001, my laboratory began to obtain transgenic rice lines (see attached Exhibit B, Lab Notebook II at page 1).
9. During approximately, June 2001 to May 2002, two generations for transgenic rice lines were analyzed for disease resistance and abiotic stress tolerance (see attached Exhibit B, Lab Notebook II at pages 2-4).
10. Prior to studies in my laboratory, no one in the field was aware that rice MAPK5 gene, its protein and enzyme activity were induced by drought, salt and low temperature and capable of rendering abiotic stress tolerance.

I certify that the foregoing statements made by me are true. I am aware that if any of the foregoing statements made by me is willfully false, I am subject to punishment.

Date: 9/25/2007



Yinong Yang Ph.D.

Department Plant Pathology
Subject Rice Defense gene Screening and Identity
Name Lizhong Xiong (NTI)
Address R. APC 215
National® Brand G-9-10 - 2001-2

Computation Notebook

11 3/4" x 9 1/4", 4 x 4 Quad., 75 Sheets

43-648



0 73333 43648 8

I

AVERY
DENNISON
OFFICE PRODUCTS
Chicopee, MA 01022

May 3 Rice seeds (Drew) planting
53g → 60 plots plots

May 4 Blots probe

N5-1 N6-1
Minwoo's chemical
Induced (including CH)
seedlings

BTH10

N5-2 N6-2
Minwoo's blots
Suspension cell

JA60

May 8-9 Culturing & Min:prep / sequencing
↓ all $A_{260}/280 \geq 1.8$ but ≤ 1.95

| | | | | | | | | |
|----|------------------|------------|----|------------------|------|----|------------------|------|
| 1 | 2A ₁₀ | 0.32 μg/ml | 11 | 2D ₁₀ | 0.29 | 21 | 2F ₁₁ | 0.34 |
| 2 | 2A ₁₂ | 0.34 | 12 | 2E ₂ | 0.31 | 22 | 2G ₁ | 0.32 |
| 3 | 2B ₁ | 0.23 | 13 | 2E ₃ | 0.30 | 23 | 2G ₅ | 0.37 |
| 4 | 2B ₇ | 0.36 | 14 | 2E ₄ | 0.5 | 24 | 2G ₆ | 0.38 |
| 5 | 2C ₁ | 0.30 | 15 | 2E ₇ | 0.41 | | | |
| 6 | 2C ₃ | 0.46 | 16 | 2E ₈ | 0.42 | | | |
| 7 | 2C ₄ | 0.38 | 17 | 2E ₁₁ | 0.37 | | | |
| 8 | 2C ₁₂ | 0.36 | 18 | 2F ₇ | 0.40 | | | |
| 9 | 2D ₂ | 0.46 | 19 | 2F ₈ | 0.29 | | | |
| 10 | 2D ₇ | 0.38 | 20 | 2F ₁₀ | 0.41 | | | |

Blast Result of JBC sequence

| SX# | Inducible data | | | Possible genes based on homology (BLASTX) |
|------|----------------|-----|-----|--|
| | Blast | BTH | JA | |
| 2A2 | | - | + | No homology |
| 2A3 | - | + | ++ | Putative Beta-ketoacyl-CoA synthase |
| 2A4 | - | + | ++ | Low homology (9E-5) with an unknown protein from Arabidopsis |
| 2A8 | + | + | ++ | No homology |
| 2A10 | - | +/- | + | No homology |
| 2A12 | - | +/- | ++ | gb AAF21081.1 AC013258_19 (AC013258) unknown protein [Arabidopsis thaliana] |
| 2B1 | - | +/- | + | No homology |
| 2B7 | - | +/- | + | 1. hypothetical protein from Arabidopsis (5E-38) 2. cytokinin oxidase-like protein (Arabidopsis) (7E-24) |
| 2B8 | - | +/- | + | hypothetical protein from Arabidopsis (5E-18) |
| 2B9 | - | - | ++ | No homology |
| 2C1 | + | - | - | No homology |
| 2C3 | ++ | + | - | RUBISCO activase |
| 2C4 | + | ⊕- | + | =2F8 |
| 2C12 | ++ | + | + | MAP kinase (high homology one from maize) |
| 2D2 | ++ | ++ | - | (AC016661) Putative ankyrin (arabidopsis) |
| 2D7 | + | - | + | (S39045) Zinc-finger protein from wheat (WZF1) Minnow |
| 2D10 | +/- | - | + | Hypothetical protein from Arabidopsis (4E-6), 24/32 (75%) |
| 2E2 | +/- | + | | (Z99707) MAP3K-like protein kinase from Arabidopsis |
| 2E3 | - | - | +/- | Not sequenced |
| 2E4 | +++ | - | ++ | No homology |
| 2E7 | R | +++ | ++ | Low homology: hypothetical protein from Arabidopsis |
| 2E8 | - | - | + | No homology |
| 2E11 | - | + | ++ | NAD-malate dehydrogenase |
| 2F6 | ++ | +/- | + | Oryza sativa mRNA for osNAC6 protein (E-155) |
| 2F7 | ⊕- | + | ++ | No homology |
| 2F8 | ++ | - | ++ | Beta-ketoacyl-CoA synthase |
| 2F10 | R | - | + | 1. An unknown protein from Arabidopsis 2. Ca+2-binding EF hand protein from soybean 3. ABA induced protein from rice |
| 2F11 | + | + | ++ | = 2A12 |
| 2G1 | ++ | +/- | ++ | No homology |
| 2G5 | R | - | - | Chlorophyll A/B binding protein |
| 2G6 | ⊕ | - | ++ | (AF225703) RSH2: Arabidopsis Rel/SpoT homology |

Add

SX3A4

SX2B7

SX1 F1

2D8

for delete redundancy

Aug 15. MAP Kinase (2C₁₂) Screening again.

Some (in 10) weeks signal dots → Continue

Aug 15 : Northern

ABJS 1

HW 1

blast 7 #

Southern 2 *

L34SP (specific probe obtained by PCR)

ABJS 2

HW 2

blast 9 *

southern 4

L68SP (specific probe)

PhosphoImager's scan: nothing bands remained!

→ ? Washing problem

→ ? Blots problem: (too old blots)

Aug 21

* Library screening with L80 (partial cDNA insert, 1 kb or so)
probe DNA was checked by gel

* Northern

ABJS 1

HW 1

blast 6 #

Southern 2

L34SP

ABJS 2

HW 2

Blast 10 (1st time use)

Southern 1 (1st time use)

L68SP

9/30

↓ Successfully excised all phage mid into plasmid.
 XLOLR cell: New tube grown in LB!

10/2

Absolutely fresh cells to be used.

$$M_{11-1} \stackrel{?}{=} [M_{11-2} = M_{11-3} = M_{11-5} = M_1] = 1.4 \text{ kb}$$

$$M_2 \stackrel{?}{=} M_8 = 1.6 \text{ kb}$$

$$M_3 2.2 \text{ kb}$$

$$M_4 0.8 \text{ kb}$$

$$M_5 \stackrel{?}{=} M_6 = 1.3 \text{ kb}$$

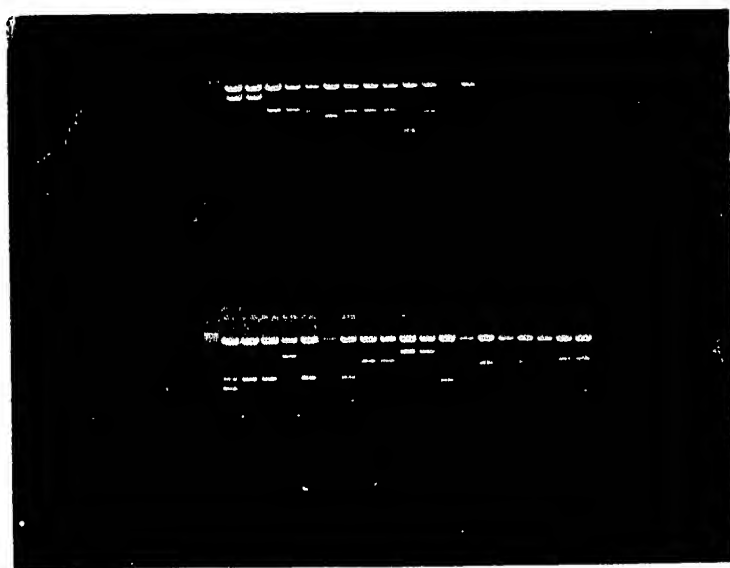
$$R_1 \stackrel{?}{=} 1.4 \text{ kb}$$

$$R_2 \stackrel{?}{=} 1.4 \text{ kb}$$

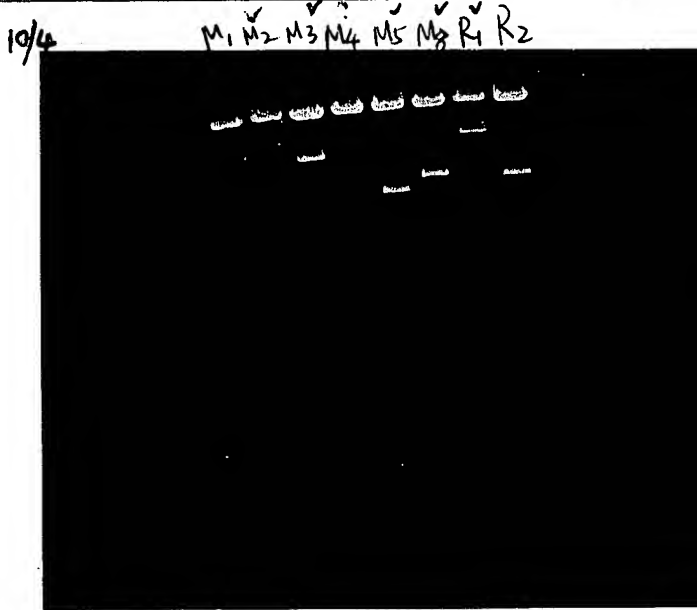
$$R_{41-1} \neq -2 \neq [-3 = -4 = -5]$$

()
maybe true $\rightarrow 1.5 \text{ kb}$

$$R_{51-1} \neq -2 \neq -3 = -4$$



| Min: Prep. | Final Cone 0.16 mg/fu) | |
|------------|---------------------------|------|
| M1 | | 1.85 |
| M2 | 0.1 | 2.0 |
| M3 | 0.36 | 1.62 |
| M4 | 0.24 | 1.84 |
| M5 | 0.25 | 1.7 |
| M8 | 0.22 | 1.8 |
| R1 | 0.11 | 0.2 |
| R2 | 0.26 | 1.8 |



10/5 Send 6 samples for sequencing < To Little Rock >

- | | | | |
|----------|-------------|------------|--|
| No. 6 M2 | = M8 | ? | 2 C ₁₂ (need further sequence or digestion) |
| No. 1 M3 | — Wrong! | → 18S RNA | |
| 2 M5 | Partial ? | = M2 or M8 | |
| 3 M8 | = M2 | | |
| 4 R1 | full-length | ? | L80 |

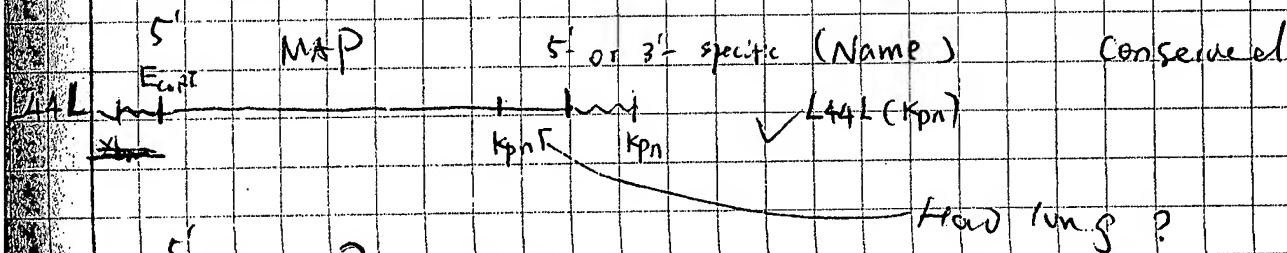
* RNP gel / RNA blotting II for Logis (CC) Test

CCBT 1 - CCBT 3

E₀, E₁, E₃, E₂₄, C₁, C₂, C₃, C₄, C₅, C₆, B₅, B₆, B₇, A₁, A₂, A₃, A₄, V₀, V₁, V₂, V₃, V₄

0:30
Pst I
Apa I

Generating Gene-specific probe (for Northern) or Conserved probe [screening homologs]



M₂ Sac I Sac I 430 \Rightarrow M₂(Sac I) = 430

(note: 2C₁₂ covers domain X, XI, so it is not gene-specific)

L68 Pst I ~~Hinf I~~ ~~Pst I~~ (620+) \Rightarrow L68 (Pst+) 620

? ✓ L68(Pst+T7) product doesn't work well

L34 ~~Hinf I~~ ~~LysP~~ \Rightarrow L34(SP+T7) (works!)

✓ PCR: L34 SP+T7 (works!)

Weekend

o Trial proposal for NOVARTIS Corporation

Jan 10

(1) Northern blotting

ABA - BTI - JA

(3x7: 0, 1/2, 1, 2, 4, 6, 12 hr.)

SA - wounding - Avir - vir

7 7 5 (0, 1/2, 1, 2, 4 days)

2 sets

Blotter name : All-in-One 1[#], 2[#]1[#] 150 ng / μ l x 400 μ l2[#] 600 ng / μ l x 500 μ l

(2) Southern blotting

3 μ g digested by Eco RI, Hind III

(perfect digestion)

New DNA from Drew using CTAB method

SEH-5, -6, -7, -8

Repeat

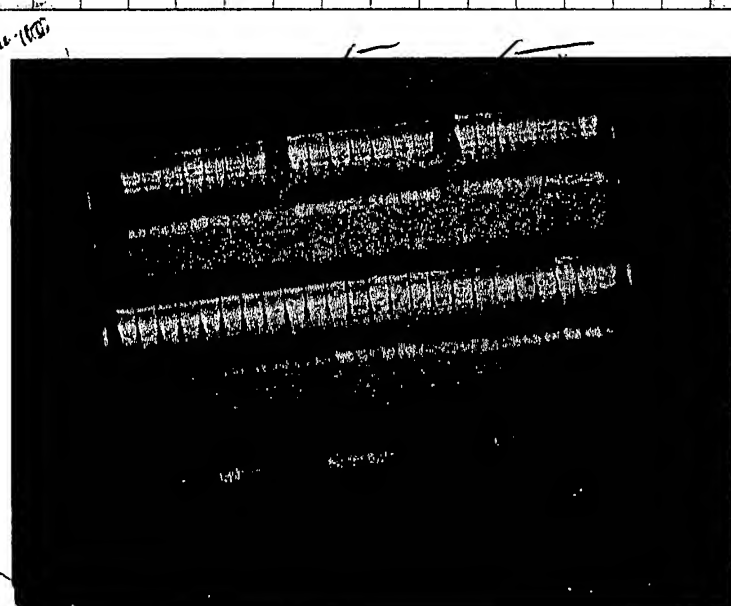
(3) Fusion construct \rightarrow ligation \rightarrow transformation

people repeat from ligation

X

(see Jan 3 for detail)

(1) Picture attached:



Department Plant Pathology
Subject Rice Defense gene Characterization
Name Lizhang Xiong
Address Rose APC 215
National Brand 2001.3

Computation Notebook

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43-648



0 73333 43648 8

II.

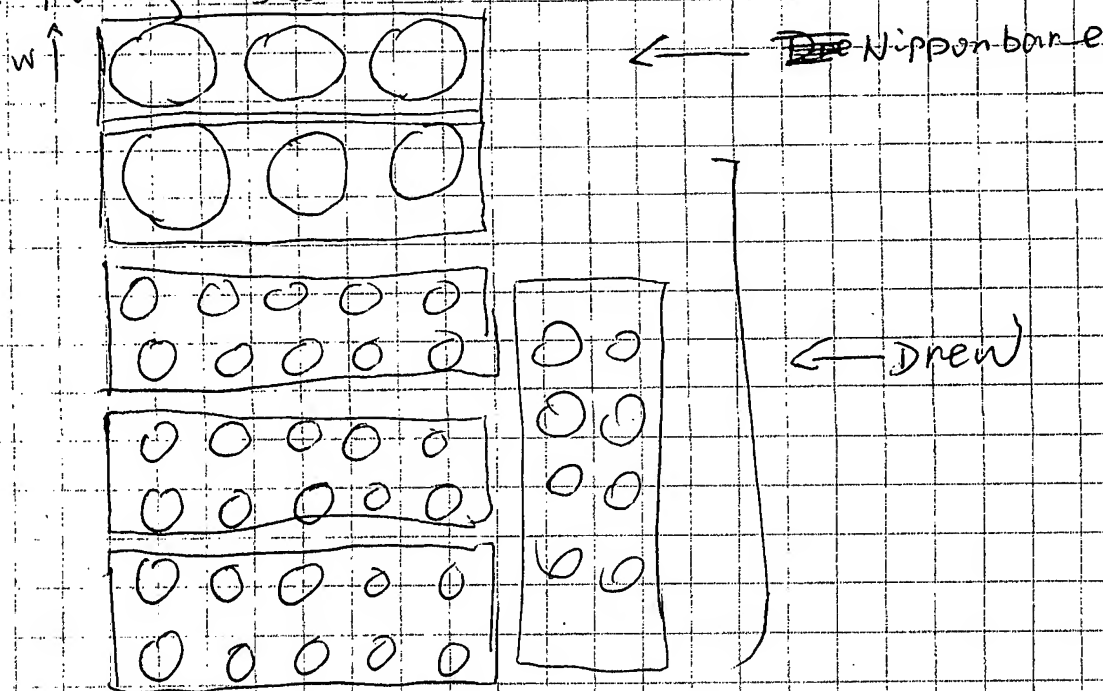


Office Products
Chicopee, MA 01022

5.18 - Summary of Transformation efficiency.

| Construct | Resistant calli | Total | Shoots obtained | Total |
|--------------------------------|------------------------------------|---------|-----------------|-------|
| R F ₂ -N | 18/37 12/41 | 84/276 | / | / |
| G ₂ -N | 24/29 22/32 17/26 | 119/301 | | |
| H H ₂ -N | 26/41 28/42 | 107/244 | | |
| H ₂ -D | 3/46 1/42 | 7/263 | | |
| C ₃ -D | 6/52 2/47 | 9/312 | | |
| C ₃ -N | (?) 2/36 0/- 4/51 | 7/282 | | |
| G ₂ -HJ | 1/42 2/40 2/38 0/- 0/- | 9/317 | | |

5/12 * Planting Seeds



Purpose : Stress Test for M_2 * large pots are control
for transgenic line

1* Salt 150-200 mM NaCl

Root \downarrow leaf. 0. ~~0.5%~~. 1. 3. 6. 10 24. 72 hr. 7d.

2* Cold $28^\circ \rightarrow 4^\circ C$:

0 3h 6h 12h 24h

or $28^\circ \rightarrow 24^\circ$ for 24h $\rightarrow 28^\circ$.

0' 3h' 6'h 12'h 24h

3* Drought

Stop water supply (wet soil)

0 day 1 d 2. 3 d 4 d

\leftarrow water content

4* Senescence (chlorophyll content?)

* Sampling : small scale in 1.5 ml tube (RNA)

medium scale in 15ml tube (protein)

Order Mycor
membrane!

D. TRIAL Western / plant protein

- ABA-induced. Wounding induced. Blast fungus induced.
- Extracted w/ Lab protocol for tobacco

E. TRANSGENIC "F₂" (M₂ - DsRNA_i)

2 Stages Experiment

STAGE I : COPY NO. (Southern) X and enzyme screening all 40 lines
 EXPRESSION OF DsRNA_i | 1st Hybridized w/ sequ. on DsRNA_i
 | 2nd --- w/ sequ. not on DsRNA_i

STAGE II : Matured plants (w/ 1 copy and expressed DsRNA_i)

* leaf segment → blast fungus (Dot inoculation)
 (Note: not 18/1, ask Min for fungus)

* intact leaf on plant → spray ABA.

other treatment using leaf segment if possible

* phenotype recording for all lines (all constructs).

Only lines showing that M₂ is inhibited to be induced
 carry on to T₁ generation.

F.

TRANSGENIC "F₂"-N / H₂-D_i

STAGE I : Same as "F₂"

STAGE II : Same as "F₂" (Focus on blast fungus)

Expected lines: Enhanced Resistance.

G. TRANSGENIC Line "G₂" - L₄₄₁ DsRNA_i

STAGE I : Same as in E. except:

Sampling for ABA at both 8 AM. 9 PM

endogenous species

6.7 1. Transfer seedlings (H_2^+) (C_3-NH)

2. Sampling: Cold-RC -24hr.
Salt 48h. (leaf & root) [AM]

Drought PM 3:00 (2 day)
plus
 $F_{2-1} = 22$

3. Extract RNA for all samples, Conc. not determined.

4. PCR for M_2 -deletion / splicing

Drew, Drew, plasmid M_2 , plasmid M_1 , H₂O

primer: RTM₂F RTM₂R (product length of M_2 should be 1.0)

Taq: Home made (0.5 μl) in 50 μl vol
added after temp. reaches 95°C

6.8 1. CK PCR

2. Transfer E.6-HJ (only one) Resistant callus to Regeneration medium

3. Salt 3d.
drought 3d. | Sample →

4. prepare talk in Mon. (SERK)